[54] RETENTION AND DRAINAGE AID FOR PAPERMAKING

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[63] Related U.S. Application Data

[51] Int. Cl. D21H 3/78
[52] U.S. Cl. 162/168.3; 162/175; 162/178; 162/181.6
[58] Field of Search 162/175, 178, 168.3, 162/181.6

[56] References Cited
U.S. PATENT DOCUMENTS
2,217,466 10/1940 Bayliss 210/23
2,244,325 6/1941 Bird 252/313
2,918,399 12/1959 Eichmeier 162/181.6
3,224,927 12/1965 Brown et al. 162/155
3,253,978 5/1966 Bodendorf et al. 162/152
4,213,950 7/1980 Mahler 423/329
4,388,150 6/1983 Sunden et al. 162/175

FOREIGN PATENT DOCUMENTS
8600100 1/1986 World Int. Prop. O. 162/181.6
8605826 10/1986 World Int. Prop. O. 162/181.6

[57] ABSTRACT
An improvement in a papermaking process in which an aqueous paper furnish containing cellulosic pulp, and optionally also mineral fillers is formed and dried, the improvement being the addition of a drainage and retention aid comprising a water soluble alkali metal polyaluminosilicate microgels formed from the reaction of polyisilicic acid and an alkali metal aluminate, the polyaluminosilicate having an alumina/silica mole ratio greater than about 1/100, together with a cationic polymer selected from the group consisting of cationic starch, cationic guar and cationic polyacrylamide.

7 Claims, No Drawings
RETENTION AND DRAINAGE AID FOR PAPERMAKING

This application is a continuation-in-part of application Ser. No. 07/143,350 filed 01/13/88 abandoned.

FIELD OF INVENTION

This invention relates to papermaking. More specifically, it relates to a method whereby a suspension of pulp and inorganic filler in water is spread over a wire or net and water is removed to form a fiber web or sheet. Even more specifically, the invention relates to the addition of water soluble anionic polyaluminosilicates microgels together with an organic cationic polymer to the pulp and filler suspension. These additives affect a flocculation of the fiber and filler fines such that during the subsequent water removal step, the ease of water removal and the retention of fines is increased thereby improving both the productivity and yield of the papermaking process.

BACKGROUND AND SUMMARY OF INVENTION

Many additive systems for improving wet-end drainage and retention have been disclosed in the prior art including those employing combinations of colloidal silica and organic polymers. Such systems are among the most efficient now in use but they are also among the most expensive and there is a continuing need to improve additive performance while reducing additive cost. Consequently, it is a primary object of this invention to provide a method whereby additive cost can be significantly reduced while at the same time increasing additive performance.

This invention employs as a retention and drainage aid a water soluble polyaluminosilicate microgel formed by the reaction of polysilicic acid with an aluminum salt, preferably an alkali metal aluminate. They consist of aggregates of very small particles having a high surface area, typically about 1000 meters²/gram (m²/g) or greater and an alumina/silica mole ratio or content greater than about 1/100 and preferably between about 1/25 and 1. Their physical structure is believed to form particle chains and three dimensional networks or microgels.

The water soluble polyaluminosilicate microgels and a process for making them are taught in co-pending U.S. Application to John Derek Rushmere CH-1554A, a Continuation-in-Part of CH-1554, both of which are incorporated herein by reference.

The polyaluminosilicates thus formed provide improved operating benefits over the aluminated colloidal silicas of the prior art in papermaking. Such prior art commercial aluminated colloidal silicas used in papermaking consist of larger, non-aggregated particles with a surface area of about 500-550 m²/g, a surface acidity of 0.66 milliequivalent per gram (meq/g) or less, and an alumina/silica mole content of about 1/60.

It is known that amorphous water insoluble polyaluminosilicates can be formed by the reaction of alkali metal polysilicates with alkali metal aluminates. Such polyaluminosilicates or synthetic zeolites have found use as catalysts, catalyst supports and ion exchange materials. Also, it is known that the particles in colloidal silica sois can be surface aluminated by aluminate ions to form a coating of polyaluminosilicate as disclosed in the book "The Chemistry of Silica" by Ralph K. Iler, John Wiley & Sons, NY, 1979, pp. 407-410.

U.S. Pat. No. 4,213,950 discloses an improved process for the preparation of amorphous, water insoluble polyaluminosilicates by the reaction of alkali metal aluminates with aqueous polysilicic acid at pH 2-4. The disclosure stresses the use of true solutions of polysilicic acid not appreciably crosslinked and distinguished from colloidal solutions, suspensions, dispersions and gels.

The new water soluble polyaluminosilicate microgels employed in this invention have unique properties and characteristics. They are formed over a wide pH range of 2-10.5 by the reaction of aqueous solutions of partially gelled polysilicic acid and an aqueous solution of an aluminum salt, preferably an alkali metal aluminate, followed by dilution of the reaction mix before gelation has occurred in order to stabilize the polyaluminosilicate microgels in an active form. Alternatively, the water soluble polyaluminosilicate microgels may be produced by dilution of the polysilicic acid stock before mixing with the alkali metal aluminates. The water soluble polyaluminosilicates so produced are distinct from the amorphous polyaluminosilicates and polyaluminosilicate coated colloidal silicas of the prior art in that they have a very high surface area, typically 1000 meter²/gram (m²/g) or greater and surprisingly a very high surface acidity, typically 1 meq/g or greater. The alumina/silica mole ratio or content is generally greater than about 1/100 and preferably between about 1/25 and 1. Their physical structure is believed to consist essentially of aggregates of very small particles of silica, surface aluminated, formed into chains and crosslinked into three-dimensional networks or microgels. Some colloidal silica and colloidal alumina particles may be present with the polyaluminosilicate microgels.

The water soluble polyaluminosilicates microgels used in this invention are believed to derive their structure from the polysilicic acid stock formed initially by an appropriate deionization or acidification of a dilute alkali metal polysilicate, for example Na₂O·3.25SiO₂. Such polysilicic acid stock, also known as "active silica" consists, according to Iler in the above cited text, p. 174 and 301-303, of very small 1-2 nanometer (nm) primary particles which are aggregated into chains and three dimensional networks or microgels. Such networks, when converted to aluminosilicates by reaction with sodium aluminate exhibit a considerably greater efficiency in flocculating fiber and filler fines than larger non-aggregated aluminated silica particles particularly when employed with a cationic polymer, such as cationic starch, cationic guar or cationic polysacrylamide. The greater efficiency in flocculation is believed to result from both the increased effectiveness of the microgel structure in locking together or bridging pulp and filler fines and also from the high surface acidity more effectively completing charge neutralization reaction with the cationic components.

The water soluble polyaluminosilicates have a wide range of application to different papermaking stocks including those containing bleached kraft pulp, groundwood pulp and thermomechanical pulp. They may also be used for the clarification of white waters and the recovery of pulp and filler components. They function well under both acid and alkaline papermaking conditions, that is, over a pH range of about 2-10.

U.S. Pat. No. 2,217,466 describes the early use of polysilicic acid or active silica as a coagulant aid in the treatment of raw water. The article "Activated Silica, a
New Chemical Engineering Tool" by Merrill and Bolton, Chem. Eng. Progress 1 (1947), 27, summarizes the development and application of anionic active silica and mentions its use as a coagulant for paper mill white water and as a retention aid for fiber and filler fines when added to the head box of a paper machine. No mention is made of the co-use of anionic active silica together with cationic polymers.

U.S. Pat. No. 3,224,927 and U.S. Pat. No. 3,253,978 disclose the co-use of cationic starch together with anionic colloidal silica as a binding agent for inorganic fibers in refractory fiber bonding applications. The quantities of colloidal silica used are considerably larger than in papermaking applications, that is, 10–20 weight percent (% wt.) of the product for fiber bonding versus about 1 wt. % of the product for paper applications. Also, in fiber binding, conditions leading to flocculations are to be avoided whereas in papermaking, flocculation is a desired result of the additions.

U.S. Pat. No. 4,388,150 discloses a binder composition comprising colloidal silica acid and cationic starch for addition to papermaking stock to improve retention of stock components or for addition to the white water to reduce pollution problems and to recover stock component values.

International Patent Publication WO86/00100 extends the application of colloidal silicas in papermaking to more acid conditions by describing the co-use of aluminated colloidal silica with cationic starches and cationic guar. Alumination provides stronger acid sites on the surface of the colloidal silica. As a consequence, anionic charge is maintained well into the acid range. The preferred compositions are those containing non-aggregated silica particles of relatively large 5–6 nm diameter, surface area of 500 m²/g and an alumina-/silica mole content of about 1/60.

International Patent Publication WO86/05826 describes the co-use of the above aluminated colloidal silica and cationic polycrylamides in papermaking.

DETAILED DESCRIPTION OF THE INVENTION

Preparation of the polyaluminosilicates used in this invention require the initial preparation of polysilicic acid microgels otherwise known as active silica. Methods for the preparation of active silica are well described in the book "Soluble Silicates," Vol. II, by James G. Vail and published by Reinhold Publishing Co., NY, 1960. In general, the methods all involve the partial acidification usually to about pH 8–9 of a dilute solution of alkali metal silicate such as sodium polysilicate Na₂O.3.2SiO₂. Acidification has been achieved using mineral acids, acid exchange resins, acid salts and acid gases. The use of some neutral salts as activators has also been described.

For the purpose of practicing the present invention, acid deionization of a dilute solution of sodium polysilicate, is preferred although the other methods of activation reported in the literature may also be used. Iler, in the above stated text at page 288, teaches that solutions containing up to 12 wt.% SiO₂ can be used in the formation of polysilicic acid, the higher percentages requiring rigorous, tightly controlled operating conditions. While the full range can be used in the practice of this invention, SiO₂ concentrations in the range of 0.1–6 wt.% is preferred. Acidification using any strong acid exchange resin known in the art, such as disclosed in U.S. Pat. No. 2,244,325, is preferred since it effectively removes the unwanted sodium value of the sodium silicate. If this sodium value is not removed and sulfuric acid, say, is used for the acidification considerable quantities of sodium sulfate are generated in the product. This sodium sulfate can be burdensome in maintaining both pollution and corrosion control standards.

The deionization is preferably conducted into the acid range of pH 2.5–5 although the higher pH ranges of 5–10.5 may also be employed particularly if higher sodium ion concentration can be tolerated. In the pH 2.5–5 range, the polysilicic acid is metastable and conditions are favorable for aggregation of the very small, high-surface-area particles into the desired chain and three dimensional networks described earlier.

The surface area of the polysilicic acids so formed generally exceeds about 1000 m²/g, typically ranging from about 1000 m²/g to 1300 m²/g, most often about 1100 m²/g. All have been found to be effective for the formation of polyaluminosilicates.

Lower SiO₂ concentrations are preferred, particularly in the preferred acid range of pH 2.5 to 5. The metastability of the polysilicic acid so formed has been found to vary with the silica concentration and method of preparation. For example, at 3 wt. % SiO₂ when prepared by batch-deionization the stability at ambient temperatures is less than a day before gelation occurs. When the polysilicic acid is formed by column-deionization, stability at ambient temperatures of greater than one day can be achieved even at 6 wt.% SiO₂. At 1 wt. % SiO₂, however, stability at ambient temperatures is excellent as measured by only small losses in surface area and no visible signs of increased viscosity or gelation over a period of three to four weeks. Further, at 1 wt. % SiO₂ concentration, surface area was found to decrease only slowly. One product with an initial surface area of 990 m²/g (as measured by the titration method of G. W. Sears, Anal. Chem. 28 (1956), 1981), decreased in surface area by only 15% over a period of a month. It was also still an effective starting material for forming polyaluminosilicates.

While aging is not essential, it has been found that generally the suitability of polysilicic acid as a precursor for the polyaluminosilicates improves with aging so long as the time of aging is less than the time it takes for the polysilicic acid to gel. That is, polyaluminosilicates prepared from a 1 wt.% polysilicic acid (polysilicic acid containing 1 wt % SiO₂), for example, that has been aged for 24 hours are frequently more effective flocculation agents than polyaluminosilicates from the same polysilicic acid when freshly prepared. The aging period has allowed time for more particle chain and three dimensional network formation.

It is important to stress the need for three dimensional network or microgel formation in the polysilicic acid stock used. While the formation of a total gel as evidenced by highly increased viscosity and water insolubility is to be avoided, the formation of the microgel is extremely important. The microgel or three dimensional network formation represents the initial stages of the gelation process before any significant increase in viscosity has occurred. Microgel formation is a function of time, silica concentration, pH and the presence of neutral salts, and significant differences can be observed in the performance of polysilicic acid formed by different modes of deionization. For example, if the deionization of a 1 wt.% SiO₂ solution, as sodium polysilicate (Na₂O.3.2SiO₂) is conducted rapidly, that is in a batch mode with a large excess of ion-exchange resin, the
polymeric acid product is likely to have little three dimensional network or microgel formation and will be less effective as a stock for polyaluminosilicate formation until it has aged. On the other hand, if the deionization is conducted slowly with successive small additions of ion-exchange resin and pH equilibration at each stage, the resultant polymeric acid will require no further aging to produce polyaluminosilicates showing excellent performance.

In practice a preferred mode of polymeric acid stock preparation is to acidify the more concentrated sodium poly-silicate solutions (3-6 wt.% SiO₂) to facilitate microgel formation and then to dilute to 1 wt.% SiO₂ or less to stabilize.

After the polymeric acid has been prepared it is mixed with the required amount of alkali metal aluminate to form the polyaluminosilicate having an alumina/silica content greater than about 1/100 and preferably 1/25 to 1. Any water soluble aluminate is suitable for this purpose. Sodium aluminates are the most readily available commercially and are therefore preferred. Solid sodium aluminate generally contains a slightly lower sodium/aluminum mole ratio than liquid sodium aluminate (that is, 1.1/1 for solid versus 1.25/1 for liquid). Lower sodium in the solid aluminate is advantageous in minimizing cost and sodium content of the polyaluminosilicates. offsetting this advantage is the considerable convenience of using the commercial liquid aluminate products.

Dilute solutions of aluminate are preferred. For example, a sodium aluminate solution containing about 2.5 wt. % Al₂O₃ prepared by diluting VSA 45, available from Vinings Chemical Co., Atlanta, GA, is suitable for this purpose.

The alkali metal aluminate must be added before the polymeric acid gels and preferably at a time that is less than 50% of the time it would take the polymeric acid to gel.

After formation, the polyaluminosilicates are diluted to whatever concentration the end use requires. For example, dilution preferably to the equivalence of 2.0 wt. % SiO₂ or less and more preferably to 0.5 wt. % or less is appropriate for addition to the papermaking process. As prepared, the polyaluminosilicates retain their high flocculation characteristics for about 24 hours.

Because of the metastability of the polyaluminosilicates and the polymeric acid precursor and the prohibitive cost of shipping stable, but very dilute, solutions containing about 1 wt. % silica, a preferred embodiment is to produce the polyaluminosilicate at the location of intended use.

The polyaluminosilicate made by the process of this invention is more reactive and efficient in the papermaking process than the commercial aluminated colloidal silicas that are currently used. They also are cheaper, particularly if made at the location of intended use. The user’s unit cost of silica in sodium poly-silicate (Na₂O.3.2SiO₂) is about one-tenth that of silica in commercial aluminated colloidal silicas.

In the papermaking process, cationic polymers, derived from natural and synthetic sources have been utilized together with the polyaluminosilicates. These cationic polymers include cationic starches, cationic guars and cationic polyacylamides, the application of which to papermaking has all been described in the prior art.

Generally, cationic starches are to be preferred since these have the advantages of low cost and of imparting dry strength to the paper. Where paper strength is not a primary requirement, use of the other polymers may be advantageous.

The cationic starch used may be derived from any of the common starch producing materials such as corn starch, potato starch and wheat starch, although the potato starches generally yield superior cationized products for the practice of this invention. Cationization is effected by commercial manufacturers using agents such as 3-chloro-2-hydroxypropyltrimethyammonium chloride to obtain cationic starches with degrees of nitrogen substitution varying between about 0.01 and 0.1 wt. % nitrogen. Any of these cationic starches may be used in conjunction with the polyaluminosilicates of the invention. A cationic potato starch with a nitrogen content of about 0.03 wt. % has been most frequently employed. In use, the polyaluminosilicates are employed in amounts ranging from about 0.01 to 1.0 wt. % (0.2 to 20 lb./ton) of the dry weight of the paper furnish together with cationic polymer in amounts ranging from about 0.01 to 2.0 wt. % (0.2 to 40 lb./ton) of the dry weight of the paper furnish. Higher amounts of either component may be employed but usually without a beneficial technical gain and with the penalty of increased costs. Generally preferred addition rates are about 0.05 to 0.2 wt. % (1-4 lb./ton) for the polyaluminosilicates together with 0.5 to 1.0 wt. % (10-20 lb./ton) of cationic starch and 0.025 and 0.5 wt. % (0.5 to 10 lb./ton) for the cationic guars and cationic polyacrylamides.

EXAMPLES

For the purpose of demonstrating the significant superiority of the polyaluminosilicates of the present invention over the aluminated colloidal silicas of the prior art, comparison tests have been made using the retention/drainage aid system marketed in the United States under the trade name “Compozil” (Procomp, Marietta, GA).

“Compozil” is a two-component system comprising BMB-a cationic potato starch and BMA-9-an aluminated colloidal silica. The BMA-9 product contains non-aggregated silica particles of surface area about 500 m²/g with an alumina to silica mole ratio of about 1/60 and a surface acidity of about 0.66 meq/g.

In conducting the comparisons, both Canadian Standard Freeness measurements for drainage and Britt Dynamic Drainage measurements for fines retention have been made. For both types of measurements mixing conditions and order of addition of the components have been maintained. Optimum results are usually obtained if the cationic polymer is added first to the papermaking furnish followed by the polyaluminosilicate, although the reverse order of addition can also be followed.

Mixing in all examples was conducted in the Britt Jar at an agitator speed of 800 rpm. For freeness measurements the treated furnish was then transferred to the cup of the freeness tester. The following mixing times were followed: (1) add furnish to Brit Jar and stir for 15 seconds, (2) add cationic polymer and stir for 15 seconds, (3) add polyaluminosilicate and stir for 15 seconds, and (4) drain for fines retention measurement or transfer to freeness tester for freeness measurement.

PREPARATION OF POLYALUMINOSILICATES

Commercial sodium poly-silicate (Na₂O.3.2SiO₂) was diluted with water to provide 500 grams of a solution
containing 1 wt. % SiO₂. To this was added slowly, in stages, about 100 grams of Dowex @50W-X8(H+), a strong sulfonic acid ion exchange resin in the acid form. The mixture was well stirred and the pH followed until it had reached a pH of about 3. The resin was removed from the polysilicic acid by filtration. With no aging period of the polysilicic acid solution, sufficient dilute sodium aluminate solution containing 2.5 wt. % Al₂O₃ was added to form the polyaluminosilicate of the desired Al₂O₃/SiO₂ ratio. The polyaluminosilicate was diluted to 0.5 wt. % SiO₂ or less for use in the following examples.

EXAMPLE 1—DRAINAGE COMPARISONS

In this example measurements were made of the drainage performance of various polyaluminosilicate compositions of the invention when used in combination with a commercial sample of “Compozi” cationic starch component BMB, S-190. All tests were made at a constant starch loading of 20 lb./ton. Comparison tests were also made using a commercial sample of “Compozi”-aluminated silica component BMA-9. All polyaluminosilicates used were freshly prepared. That is, just prior to the tests, fresh polysilicic acid containing 1 wt. % SiO₂ prepared by acid deionization of sodium polysilicate, Na₂O·3.2SiO₂ was mixed with the desired amount of dilute sodium aluminate (2.5 wt. % Al₂O₃) and the mixture was then diluted to 0.5 wt. % or less.

The furnish used was a fine paper furnish containing 70% bleached kraft pulp (70% hardwood, 30% softwood), 20% kaolin clay and 1% calcium carbonate. To this, 0.66 g/l of anhydrous sodium sulfate was added as electrolyte and the pH was adjusted to 4.5 by the addition of sulfuric acid. The furnish was made up at 0.5 wt. % consistency but diluted to 0.3 wt. % consistency for freeness measurements.

The results are given in Table 1, from which it may be seen that the polyaluminosilicates of the invention out-performed the commercial sample of aluminated colloidal silica (BMA-9). The more preferred polyaluminosilicates, namely those with Al₂O₃/SiO₂ mole ratios of 13/87 and 17/83 gave significantly higher drainage values even when using considerably less material. For instance, BMA-9 at a typical commercial loading of 4 lb./t gave a freeness of 385 ml whereas the 13/87 polyaluminosilicate gave an essentially equivalent freeness of 395 ml at a loading of only 1 lb./t—a fourfold reduction in material use.

EXAMPLE 2—DRAINAGE COMPARISONS

In this example measurements were made of the drainage performance of the 13/87 polyaluminosilicate when used in conjunction with various cationic starches. The polyaluminosilicate loading was held constant at 3 lb./t and the starch loading varied between 0 and 40 lb./t. A comparison was also made with the BMA-9/BMB combination of the commercial Compozi system under the same variables. The furnish used was of the same composition to that used in Example 1 and the pH was again 4.5. The starches used were:

BMB S-190—a cationic potato starch imported from Europe for “Compozi”
Stalok @400—a cationic potato starch manufactured in the U.S. by A. F. Staley Co., Decatur, IL, and
Stalok @324—a cationic waxy corn starch manufactured in the U.S. by A. F. Staley Co., Decatur, IL.

The results in Table 2 shows that the 13/87 polyaluminosilicate of the invention when used in combination with either of the cationic potato starches (BMB S-190 or Stalok @400) clearly out-performed the commercial BMA-9/BMB system. Larger drainage values were obtained at lower starch loadings—an economy in papermaking operations where dry strength is not a primary requirement. The performance of the cationic waxy corn starch (Stalok @324) was inferior as has been found to be the case generally with the lower molecular weight starches.

EXAMPLE 3—DRAINAGE COMPARISONS

In this example, drainage measurements have been made for the 13/87 polyaluminosilicate in an alkaline furnish at pH 8. The furnish was a similar composition to that used in Example 1 except that precipitated calcium carbonate replaced the clay as inorganic filler. All tests were made at a constant cationic starch loading of 20 lb./t. The starch used was BMB S-190. Comparison measurements were also made using aluminated colloidal silica of the prior art (BMA-9), simple polysilicic acid (non-aluminated) and also sodium aluminate alone.

The results are given in Table 3 and again show that the 13/87 polyaluminosilicate gives significantly improved freeness at lower loadings compared to the prior art sol. It may also be seen that the polysilicic acid alone and sodium aluminate alone (but both used in conjunction with 20 lb./t cationic starch) have no effect in improving freeness. It is their reaction product, the polyaluminosilicate of the invention, that effects improvements.

EXAMPLE 4—FINES RETENTION

In this example, measurements of fines retention were made using a Britt Dynamic Drainage Jar. The furnish used was an alkaline furnish at pH 8 of the same composition to that used in Example 3. The polyaluminosilicate used was that containing the 13/87 mole ratio of Al₂O₃/SiO₂ and comparison was again made to BMA-9 aluminated colloidal silica. Sol loading was held constant in each case at 6 lb./t and the starch loading varied between 4 and 20 lb./t. Results are in Table 4.

Using the polyaluminosilicate of the invention very significant improvements in fines retention were obtained at all starch loadings, particularly in the common commercial range of 12–20 lb./t. Compared to the prior art system, economies in paper manufacture could be obtained by the need to use less starch to maintain the same level of fines retention.

EXAMPLE 5—DRAINAGE TEST USING STONEGROUND WOOD

In order to demonstrate the wide applicability of the polyaluminosilicates to papermaking pulp systems fines measurements were made on a 0.3 wt. % furnish comprising 100% groundwood wood (aspen) under very acid conditions, pH 4.0. Stoneground wood represents the coarse end of pulp systems, whereas bleached kraft pulp represents the fine end. Stoneground wood is characterized by poor drainage (freeness) and high fines content. The results recorded in Table 5 show how increasing the amounts of 13/87 polyaluminosilicate used in conjunction with 20 lb./t cationic starch (BMB S-190) increased the freeness of the pulp system. Turbidity measurements for the white water from the freeness tests are also recorded. Decreasing turbidity is an indication of improved fines retention.
EXAMPLE 6—DRAINAGE TEST

In this example, a comparison was made of the drainage of polyaluminosilicate/cationic guar combinations versus aluminated colloidal silica/cationic guar combinations of the prior art. The polyaluminosilicate was a freshly prepared 13/97, Al₂O₃/SiO₂ mole ratio product, the aluminated silica soil was a commercial BMA-9 sample and the cationic guar was Jaguar @C-13 (Stein, Hall & Co., NY, NY). Comparisons were made using both a clay-filled furnish similar to that of Example 1 at pH 4.5 and a calcium carbonate filled furnish similar to that of Example 3 at pH 8.0. Results are given in Table 6. All tests were made at a constant guar addition of 4 lb./t (0.2 wt. %). The superiority of the polyaluminosilicate/cationic guar combinations over the prior art aluminated silica sol/cationic guar combinations is clearly demonstrated.

EXAMPLE 7—DRAINAGE TESTS

In this example a comparison is made of the drainage benefits of a polyaluminosilicate/cationic polyacrylamide combination over an aluminated silica sol/cationic polyacrylamide combination of the prior art. The polyaluminosilicate was a freshly prepared 13/37 mole product, the aluminated colloidal silica was a commercial sample of BMA-9 and the cationic polyacrylamide was a sample of Hyperfloc @605 (Hychem Inc., Tampa, Fla.) with a mol wt. of about 10 million (MM) and with a cationic content of 20–30 wt. %. Table 7 lists the results obtained in a calcium carbonate filled furnish at pH 8 similar to Example 3 and shows improved drainage performance of the polyalumino-cationic polyacrylamide combination over the prior art. All tests were made at 2 lb./t (0.1 wt. %) of cationic polyacrylamide.

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### TABLE 1

<table>
<thead>
<tr>
<th>Polyaluminosilicate</th>
<th>Freeness, ml at Sol Loading of</th>
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<tr>
<td>Al₂O₃/SiO₂</td>
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<td>Mole Ratio</td>
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<tr>
<td>2/98 (BMA-9)</td>
<td>330 330 345 385 420</td>
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<td>4/96</td>
<td>330 365 374 340 —</td>
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<tr>
<td>7/93</td>
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<td>330 395 460 505 465</td>
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<td>17/93</td>
<td>330 395 475 500 —</td>
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### TABLE 2

<table>
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<th>Starch Used</th>
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<tr>
<td>Furnish</td>
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</tr>
<tr>
<td>BMB-5-190</td>
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</tr>
<tr>
<td>(Composit)</td>
<td>310</td>
</tr>
<tr>
<td>BMB-5-190</td>
<td>310</td>
</tr>
<tr>
<td>Stalok 400</td>
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<td>Stalok 324</td>
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### TABLE 3

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<tr>
<th>Sol Used</th>
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<tr>
<td>BMA-9</td>
<td>285 330 380 415 440</td>
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### TABLE 4

<table>
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<tr>
<th>Sol Type</th>
<th>Fines Retention at Cationic Starch Loading of</th>
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<tr>
<td>BMA-9</td>
<td>27 36 42 46 49 46</td>
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<tr>
<td>Polyaluminosilicate</td>
<td>27 40 60 73 74 82</td>
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### TABLE 5

<table>
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<tr>
<th>Sol Loading of</th>
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<th>Turbidity N/T/ Units</th>
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<tr>
<td>Furnish only</td>
<td>4.5 440</td>
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</tr>
<tr>
<td>BMA-9</td>
<td>4.5 530 480 490 510 530 580</td>
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<td>Polyaluminosilicate</td>
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<td>Furnish only</td>
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<tr>
<td>BMA-9</td>
<td>8.0 390 430 470 570 660 695</td>
<td></td>
</tr>
<tr>
<td>Polyaluminosilicate</td>
<td>8.0 390 430 470 570 660 695</td>
<td></td>
</tr>
</tbody>
</table>

### TABLE 6

<table>
<thead>
<tr>
<th>Sol Used</th>
<th>Freeness, ml at Sol Addition of (lb./ton)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BMA-9</td>
<td>390</td>
</tr>
<tr>
<td>Polyaluminosilicate</td>
<td>580 660 680 670</td>
</tr>
</tbody>
</table>

### TABLE 7

<table>
<thead>
<tr>
<th>Sol Used</th>
<th>Freeness, ml at Sol Loading of</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 lb./t</td>
<td>2 lb./t 4 lb./t 6 lb./t 8 lb./t</td>
</tr>
<tr>
<td>BMA-9</td>
<td>390</td>
</tr>
<tr>
<td>Polyaluminosilicate</td>
<td>390</td>
</tr>
</tbody>
</table>

I claim:

1. In a papermaking process in which an aqueous paper furnish containing cellulose pulp, and optionally also mineral fillers is formed and dried, the improvement comprising adding to said pulp from about 0.01 to about 1.0 wt. percent, based on the dry weight of the
paper furnish, of a water soluble alkali metal polyaluminosilicate microgel formed from the reaction of polysilicic acid and an alkali metal aluminate and comprising aggregates of particles in which each particle has a surface area of at least about 1000 meters\(^2\)/gram, the polyaluminosilicate microgel having an alumina/silicate mole ratio greater than about 1/100, and from about 0.01 to about 2.0 wt. percent, based on the dry weight of the paper furnish of a water soluble cationic polymer capable of flocculating fiber and filler fines.

2. The process of claim 1 in which the polyaluminosilicate has an alumina/silica mole ratio between about 1/15 and 1/3.

3. The process of claim 1 in which the polyaluminosilicate has an alumina/silica mole ratio between about 1/6 and 1/7.

4. The process of claim 1 in which the alkali metal aluminate is sodium aluminate.

5. The method of claim 4 in which the polyaluminosilicate has an alumina/silica mole ratio between about (1/15) 1/25 and 1/7.

6. The process of claim 4 in which the polyaluminosilicate has an alumina/silica mole ratio between about 1/6 and 1/7.

7. The process of claim 1 or claim 4 in which the water soluble cationic polymer is selected from the group consisting of cationic starch, cationic guar and cationic polyacrylamide.

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